WEST Search History

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DATE: Wednesday, November 30, 2005

Hide?	Set Name	Query	Hit Count
	DB=PGPB	I, USPT, USOC, EPAB, JPAB, DWPI, TDBD; PLUR = 1	YES; OP = OR
	L1	singh.in. and anthra\$	170
	L2	L1 and iota\$	3
	L3	yogendra	1740
	L4	L3 and anthra\$	156
	L5	L4 and iota\$	3
	L6	iota\$.clm. and (anthrax\$ or anthra\$).clm.	12
	L7	(khanna or hemant)in. and anthra\$	3682
	L8	L7 and iota\$	7
	L9	5,935,990.pn.	2

END OF SEARCH HISTORY

DOCUMENT-IDENTIFIER: US 20020048590 A1

TITLE: Targeting antigens to the MHC class I processing pathway with an anthrax toxin fusion protein

CLAIMS:

- 1. A vaccine capable of inducing an immune response in a mammal to a specific antigen wherein the vaccine comprises a unit dose of an <u>anthrax</u> protective antigen and said specific antigen bound to an <u>anthrax</u> protective antigen binding protein.
- 6. The vaccine of claim 1 wherein the <u>anthrax</u> protective antigen binding protein is the lethal factor of Bacillus anthracis.
- 7. The vaccine of claim 1 wherein the <u>anthrax</u> protective antigen binding protein comprises at least about the first 250 amino acid residues of the lethal factor of Bacillus <u>anthracis</u> and less than all of the amino acid residues of the lethal factor.
- 8. The vaccine of claim 1 wherein the molar ratio of protective antigen to the antigen bound to an anthrax protective antigen binding protein is greater than one.
- 9. A method of immunizing a mammal against an antigen which comprises administering a safe and effective amount of a vaccine comprising an <u>anthrax</u> protective antigen and said antigen bound to an <u>anthrax</u> protective antigen binding protein.
- 14. The method of claim 9 wherein the <u>anthrax</u> protective antigen binding protein is the lethal factor of Bacillus anthracis.
- 15. The method of claim 9 wherein the <u>anthrax</u> protective antigen binding protein comprises at least about the first 250 amino acid residues of the lethal factor of Bacillus <u>anthracis</u> and less than all of the amino acid residues of the lethal factor.
- 16. The method of claim 9 wherein the molar ratio of protective antigen to the antigen bound to an anthrax protective antigen binding protein is greater than one.
- 19. The method of claim 9 wherein the vaccine is administered in a unit dose that is between 10 to 500 nanograms of antigen bound to an <u>anthrax</u> protective antigen binding protein per kilogram of said mammal.
- 20. A method of inducing antigen presenting mammalian cells to present specific antigens on their cell membranes via the MHC class I processing pathway, comprising: i) selecting cells that can process and present specific antigens on their cell membranes via the MHC class I processing pathway; ii) contacting the cells with an <u>anthrax</u> protective antigen and said specific antigen bound to an <u>anthrax</u> protective antigen binding protein; and, iii) permitting the cells to internalize, process and present said specific antigen bound to an <u>anthrax</u> protective antigen binding protein on its cell membrane, forming a specific antigen presenting cell.
- 23. The method of claim 20 wherein the <u>anthrax</u> protective antigen binding protein is the lethal factor of Bacillus <u>anthracis</u>.
- 24. The method of claim 20 wherein the <u>anthrax</u> protective antigen binding protein comprises at least about the first 250 amino acid residues of the lethal factor of Bacillus <u>anthracis</u> and less than all of the

amino acid residues of the lethal factor.

- 25. The method of claim 20 wherein the molar ratio of protective antigen to the antigen bound to an <u>anthrax</u> protective antigen binding protein is greater than one.
- 27. A vaccine for inducing an immune response in a mammal to a specific antigen wherein the vaccine comprises a unit dose of a binary toxin protective antigen and the antigen bound to a binary toxin protective antigen binding protein wherein the binary toxin is selected from the group comprising iota toxin and anthrax toxin.
- 28. The vaccine of claim 27, wherein the binary toxin is iota toxin.

Record List Display Page 1 of 1

20030198651. 27 May 03. 23 Oct 03. Targeting antigens to the MHC class I processing pathway with an anthrax toxin fusion protein. Klimpel, Kurt, et al. 424/246.1; A61K039/07.

☐ 8. 20030198595. 17 May 02. 23 Oct 03. Use of bi-specific antibodies for pre-targeting diagnosis and therapy. Goldenberg, David M., et al. 424/1.49; 530/391.1 534/11 A61K051/00 C07K016/46.

☐ 9. 20030148409. 15 Oct 02. 07 Aug 03. Direct targeting binding proteins. Rossi, Edmund, et al. 435/7.23; 424/1.49 530/388.8 A61K051/00 G01N033/574 C07K016/30.

☐ 10. 20020048590. 09 May 01. 25 Apr 02. Targeting antigens to the MHC class I processing pathway with an anthrax toxin fusion protein. Klimpel, Kurt, et al. 424/246.1; A61K039/07.

1.51		
	-	 100
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L5: Entry 1 of 4

File: PGPB

Apr 25, 2002

DOCUMENT-IDENTIFIER: US 20020048590-AT

TITLE: Targeting antigens to the MHC class I processing pathway with an anthrax toxin fusion protein

CLAIMS:

27. A vaccine for inducing an immune response in a mammal to a specific antigen wherein the vaccine comprises a unit dose of a binary toxin protective antigen and the antigen bound to a binary toxin protective antigen binding protein wherein the binary toxin is selected from the group comprising iota toxin and anthrax toxin.

28. The vaccine of claim 27, wherein the binary toxin is iota toxin.

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L5: Entry 2 of 4

File: USPT

Apr 10, 2001

DOCUMENT-IDENTIFIER: US 6214602 B1

TITLE: Host cells for expression of clostridial toxins and proteins

CLAIMS:

3. The host cell of claim 2, wherein said clostridial proteins are selected from the group consisting of light chains of botulinal neurotoxins, heavy chains of botulinal neurotoxins, botulinal C3 protein, clostridial <u>iota toxin</u> Ia protein, and light and heavy chains of tetanus <u>toxin</u>.

AUTHORS' CORRECTIONS

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1 47 1

3051

Characterization of *Clostridium perfringens* Iota-Toxin Genes and Expression in *Escherichia coli*

SYLVIE PERELLE, MARYSE GIBERT, PATRICE BOQUET, AND MICHEL R. POPOFF

Laboratoire des Toxines Microbiennes, Institut Pasteur, 28 Rue du Dr. Roux, 75724 Paris Cedex 15, France

Volume 61, no. 12, p. 5147-5156: A DNA sequence mistake has been found at the end of the Ia gene. The five C-terminal amino acids of Ia are incorrect, and the Ia sequence is 67 amino acids longer. The Ia and Ib genes are separated by 40 noncoding nucleotides; the Ib sequence is not changed. The Ia sequence has been corrected in the EMBL data library and should appear as shown below:

```
1 MKKVNKSISV FLILYLILTS SFPSYTYAQD LQIASNYITD RAFIERPEDF
51 LKDKENAIQW EKKEAERVEK NLDTLEKEAL ELYKKDSEQI SNYSQTRQYF
101 YDYQIESNPR EKEYKNLRNA ISKNKIDKPI NVYYFESPEK FAFNKEIRTE
151 NQNEISLEKF NELKETIQDK LFKQDGFKDV SLYEPGNGDE KPTPLLIHLK
201 LPKNTGMLPY INSNDVKTLI EQDYSIKIDK IVRIVIEGKQ YIKAEASIVN
251 SLDFKDDVSK GDLWGKENYS DWSNKLTPNE LADVNDYMRG GYTAINNYLI
301 SNGPLNNPNP ELDSKVNNIE NALKLTPIPS NLIVYRRSGP QEFGLTLTSP
351 EYDFNKIENI DAFKEKWEGK VITYPNFIST SIGSVNMSAF AKRKILLRIN
401 IPKDSPGAYL SAIPGYAGEY EVLLNHGSKF KINKVDSYKD GTVTKLILDA
```

Characterization of the Structural Elements in Lipid A Required for Binding of a Recombinant Fragment of Bactericidal/Permeability-Increasing Protein rBPI₂₃

HÉLÈNE GAZZANO-SANTORO, JAMES B. PARENT, PAUL J. CONLON, HERBERT G. KASLER, CHAO-MING TSAI, DEBORAH A. LILL-ELGHANIAN, AND RAWLE I. HOLLINGSWORTH

Sepsis Research Department, XOMA Corporation, Berkeley, California 94710; Neurocrine Biosciences, San Diego, California 92121; Department of Health and Human Services, Center for Biologics Evaluation and Research, Food and Drug Administration, Bethesda, Maryland 20892; and Departments of Biochemistry and Chemistry, Michigan State University, East Lansing, Michigan 48824

Volume 63, no. 6, p. 2201-2205: We failed to cite the study by Holst et al. in which the original structure of *Escherichia coli* J5 lipid A was first reported (O. Holst, S. Müller-Loennies, B. Lindner, and H. Brade, Eur. J. Biochem. 214:695-701, 1993).

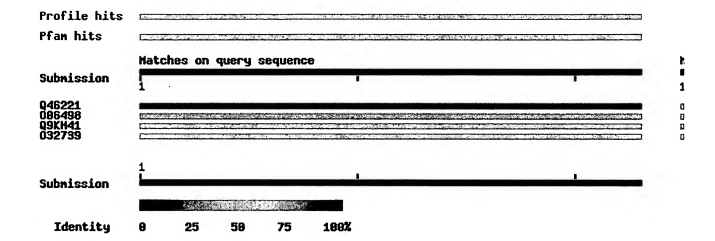
☐ tr Q9KH41 _CLODI CdtB [cdtB] [Clostridium difficile] ☐ tr O32739 CLODI ADP-ribosyltransferase [cdtB] [Clostridium diffi

Graphical overview of the alignments

Click here

to resubmit your query after masking regions matching PROSITE profiles or Pfam HMMs

(Help) (use ScanProsite for more details about PROSITE matches)



Alignments

tr Q46221 Iota toxin component Ib precursor [Clostridium Q46221 CLOPE perfringens]

Score = 73.2 bits (165), Expect = 7e-13
Identities = 23/23 (100%), Positives = 23/23 (100%)

Query: 1 DANTVGVSISAGYQNGFTGNITT 23

DANTVGVSISAGYQNGFTGNITT

Sbjct: 332 DANTVGVSISAGYQNGFTGNITT 354

Sperd

875

AA aliqn

tr 006498 Sb component [sbs] [Clostridium spiroforme] 879 AA 006498_9MOLU

align

Score = 52.8 bits (117), Expect = 9e-07 Identities = 17/23 (73%), Positives = 17/23 (73%)

Query: 1 DANTVGVSISAGYQNGFTGNITT 23

DANT GV I YQNGFTG ITT Sbjct: 336 DANTAGVAINIAYQNGFTGSITT 358

tr Q9KH41 CdtB [cdtB] [Clostridium difficile] 876 AA Q9KH41_CLODI align

Score = 49.8 bits (110), Expect = 7e-06 Identities = 15/23 (65%), Positives = 18/23 (78%)

Query: 1 DANTVGVSISAGYQNGFTGNITT 23 + NT GVS+ GYQNGFT N+TT Sbjct: 333 ESNTAGVSVNVGYQNGFTANVTT 355

tr 032739 ADP-ribosyltransferase [cdtB] [Clostridium difficile] 876 AA 032739_CLODI align

Score = 49.8 bits (110), Expect = 7e-06 Identities = 15/23 (65%), Positives = 18/23 (78%)

Query: 1 DANTVGVSISAGYQNGFTGNITT 23 + NT GVS+ GYQNGFT N+TT Sbjct: 333 ESNTAGVSVNVGYQNGFTANVTT 355

Database: EXPASY/UniProtKB

Posted date: Nov 21, 2005 2:19 PM Number of letters in database: 854,910,163 Number of sequences in database: 2,618,771

Lambda K H 0.349 0.280 1.64

Gapped

Lambda K H

0.294 0.110 0.610

Matrix: PAM30

Gap Penalties: Existence: 9, Extension: 1

Number of HSP's successfully gapped in prelim test: 0

length of query: 23

length of database: 854,910,163

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effective HSP length: 14
effective length of query: 9
effective length of database: 818,247,369
effective search space: 7364226321
effective search space used: 7364226321
T: 16
A: 40
X1: 14 ( 7.0 bits)
X2: 35 (14.8 bits)
X3: 58 (24.6 bits)
S1: 40 (22.0 bits)
S2: 62 (29.5 bits)
```

Wallclock time: 2 seconds

ExPASy Home page Site Map Search ExPASy Contact us Proteomics tools Swiss-Pro

CLUSTAL W (1.82) multiple sequence alignment

sp P13423 PAG_BACAN tr Q68GS1 Q68GS1_BACAN tr Q52NH4 Q52NH4_BACAN tr Q4ZE94 Q4ZE94_BACAN	MKKRKVLIPLMALSTILVSSTGNLEVIQAEVKQENRLLNESESSSQGLLGMEVKQENRLLNESESSSQGLLG MKKRKVLIPLMALSTILVSSTGNLEVIQAEVKQENRLLNESESSSQGLLG
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sp P13423 PAG_BACAN tr Q68GS1 Q68GS1_BACAN tr Q52NH4 Q52NH4_BACAN tr Q4ZE94 Q4ZE94_BACAN	SPEKWSTASDPYSDFEKVTGRIDKNVSPEARHPLVAAYPIVHVDMENIIL SPEKWSTASDPYSDFEKVTGRIDKNVSPEARHPLVAAYPIVHVDMENIIL SPEKWSTASDPYSDFEKVTGRIDKNVSPEARHPLVAAYPIVHVDMENIIL SPEKWSTASDPYSDFEKVTGRIDKNVSPEARHPLVAAYPIVHVDMENIIL
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sp P13423 PAG_BACAN tr Q68GS1 Q68GS1_BACAN tr Q52NH4 Q52NH4_BACAN tr Q4ZE94 Q4ZE94_BACAN	AGFSNSNSSTVAIDHSLSLAGERTWAETMGLNTADTARLNANIRYVNTGT AGFSNSNSSTVAIDHSLSLAGERTWAETMGLNTADTARLNANIRYVNTGT AGFSNSNSSTVAIDHSLSLAGERTWAETMGLNTADTARLNANIRYVNTGT AGFSNSNSSTVAIDHSLSLAGERTWAETMGLNTADTARLNANIRYVNTGT
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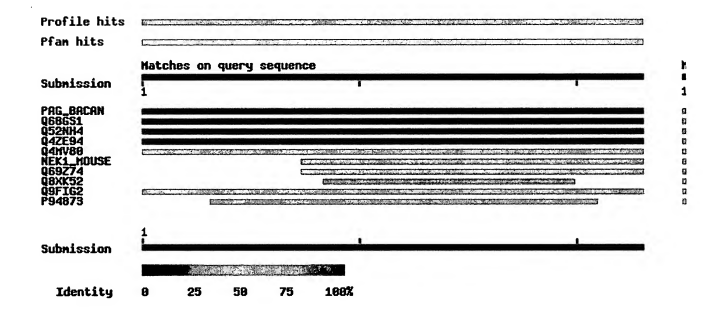
sp P13423 PAG_BACAN
tr Q4ZE94 Q4ZE94 BACAN DTGSNWSEVLPQIQETTARIIFNGKDLNLVERRIAAVNPSDPLETTKPDM ************************************
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sp P13423 PAG_BACAN TLKEALKIAFGFNEPNGNLQYQGKDITEFDFNFDQQTSQNIKNQLAELNA tr Q68GS1 Q68GS1_BACAN TLKEALKIAFGFNEPNGNLQYQGKDITEFDFNFDQQTSQNIKNQLAELNA tr Q52NH4 Q52NH4_BACAN TLKEALKIAFGFNEPNGNLQYQGKDITEFDFNFDQQTSQNIKNQLAELNA
tr Q68GS1 Q68GS1 BACAN TLKEALKIAFGFNEPNGNLQYQGKDITEFDFNFDQQTSQNIKNQLAELNA tr Q52NH4 Q52NH4 BACAN TLKEALKIAFGFNEPNGNLQYQGKDITEFDFNFDQQTSQNIKNQLAELNA
tr Q52NH4 Q52NH4_BACAN TLKEALKIAFGFNEPNGNLQYQGKDITEFDFNFDQQTSQNIKNQLAELNA
- EYLOAZROA LOAZROA - RACAN - TIKRALKIARGENEDNONLOVOGENTTERDENEDOOTSONTENDIARINA

sp P13423 PAG_BACAN TNIYTVLDKIKLNAKMNILIRDKRFHYDRNNIAVGADESVVKEAHREVIN
tr Q68GS1 Q68GS1_BACAN TNIYTVLDKIKLNAKMNILIRDKRFHYDRNNIAVGADESVVKEAHREVIN
tr Q52NH4 Q52NH4_BACAN TNIYTVLDKIKLNAKMNILIRDKRFHYDRNNIAVGADESVVKEAHREVIN
tr Q4ZE94 Q4ZE94_BACAN TNIYTVLDKIKLNAKMNILIRDKRFHYDRNNIAVGADESVVKEAHREVIN
sp P13423 PAG_BACAN SSTEGLLLNIDKDIRKILSGYIVEIEDTEGLKEVINDRYDMLNISSLRQD
tr Q68GS1 Q68GS1_BACAN SSTEGLLLNIDKDIRKILSGYIVEIEDTEGLKEVINDRYDMLNISSLRQD
tr Q52NH4 Q52NH4_BACAN SSTEGLLNIDKDIRKILSGYIVEIEDTEGLKEVINDRYDMLNISSLRQD
tr Q4ZE94 Q4ZE94_BACAN SSTEGLLNIDKDIRKILSGYIVEIEDTEGLKEVINDRYDMLNISSLRQD
sp P13423 PAG_BACAN GKTFIDFKKYNDKLPLYISNPNYKVNVYAVTKENTIINPSENGDTSTNGI
tr Q68GS1 Q68GS1_BACAN GKTFIDFKKYNDKLPLYISNPNYKVNVYAVTKENTIINPSENGDTSTNGI
tr Q52NH4 Q52NH4_BACAN GKTFIDFKKYNDKLPLYISNPNYKVNVYAVTKENTIINPSENGDTSTNGI
tr Q4ZE94 Q4ZE94_BACAN GKTFIDFKKYNDKLPLYISNPNYKVNVYAVTKENTIINPSENGDTSTNGI
sp P13423 PAG_BACAN KKILIFSKKGYEIG
tr Q68GS1 Q68GS1_BACAN KKILIFSKKGYEIG
tr Q52NH4 Q52NH4_BACAN KKILIFSKKGYEIG
tr Q4ZE94 Q4ZE94_BACAN KKILIFSKKGYEIG

Graphical overview of the alignments

Clickhere to resubmit your query after masking regions matching PROSITE profiles or Pfam HMMs

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Alignments

sp P13423 Protective antigen precursor (PA) (PA-83) (PA83) (Anthrax 764
PAG_BACAN toxins AA
translocating protein) [Contains: Protective antigen PA-20 (PA20); Protective antigen PA-63 (PA63)] [pagA]
[Bacillus anthracis]

Score = 74.4 bits (168), Expect = 3e-13
Identities = 23/23 (100%), Positives = 23/23 (100%)

Query: 1 THTSEVHGNAEVHASFFDIGGSV 23

THTSEVHGNAEVHASFFDIGGSV

Sbjct: 327 THTSEVHGNAEVHASFFDIGGSV 349

tr Q68GS1 Protective antigen [Bacillus anthracis] 736 AA

Q68GS1_BACAN

align

Score = 74.4 bits (168), Expect = 3e-13

Identities = 23/23 (100%), Positives = 23/23 (100%)

Query: 1 THTSEVHGNAEVHASFFDIGGSV 23

THTSEVHGNAEVHASFFDIGGSV

Sbjct: 299 THTSEVHGNAEVHASFFDIGGSV 321

tr Q52NH4 Protective antigen [pag] [Bacillus anthracis] 764 AA

Q52NH4_BACAN

align

Score = 74.4 bits (168), Expect = 3e-13

Identities = 23/23 (100%), Positives = 23/23 (100%)

Query: 1 THTSEVHGNAEVHASFFDIGGSV 23

THTSEVHGNAEVHASFFDIGGSV

Sbjct: 327 THTSEVHGNAEVHASFFDIGGSV 349

tr Q4ZE94 Protective antigen (Fragment) [pa] [Bacillus anthracis] 561 AA

Q4ZE94_BACAN

align

Score = 74.4 bits (168), Expect = 3e-13

Identities = 23/23 (100%), Positives = 23/23 (100%)

Query: 1 THTSEVHGNAEVHASFFDIGGSV 23

THTSEVHGNAEVHASFFDIGGSV

Sbjct: 124 THTSEVHGNAEVHASFFDIGGSV 146

CLUSTAL W (1.81) multiple sequence alignment

unk VIRT2310 Blast_submission tr Q46221 Q46221_CLOPE	MNIQIKNVFSFLTLTAMISQTLSYNVYAQTTTQNDTNQKEEITNENTLS
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unk VIRT2310 Blast_submission tr Q46221 Q46221_CLOPE	NITT NITTSYSHTTDNSTAVQDSNGESWNTGLSINKGESAYINANVRYYNTGT ****
unk VIRT2310 Blast_submission tr Q46221 Q46221_CLOPE	PMYKVTPTTNLVLDGETLATIKAQDNQIGNNLSPNETYPKKGLSPLALN
unk VIRT2310 Blast_submission tr Q46221 Q46221_CLOPE	MDQFNARLIPINYDQLKKLDSGKQIKLETTQVSGNYGTKNSQGQIITEG
unk VIRT2310 Blast_submission tr Q46221 Q46221_CLOPE	SWSNYISQIDSVSASIILDTGSQTFERRVAAKEQGNPEDKTPEITIGEA
unk VIRT2310 Blast_submission tr Q46221 Q46221_CLOPE	KKAFSATKNGELLYFNGIPIDESCVELIFDDNTSEIIKEQLKYLDDKKI
unk VIRT2310 Blast_submission tr Q46221 Q46221_CLOPE	NVKLERGMNILIKVPSYFTNFDEYNNFPASWSNIDTKNQDGLQSVANKL
unk VIRT2310 Blast_submission tr Q46221 Q46221_CLOPE	GETKIIIPMSKLKPYKRYVFSGYSKDPSTSNSITVNIKSKEQKTDYLVP
unk VIRT2310 Blast_submission	

tr Q46221 Q46221_CLOPE	KDYTKFSYEFETTGKDSSDIEITLTSSGVIFLDNLSITELNSTPEILKE
unk VIRT2310 Blast_submission tr Q46221 Q46221_CLOPE	EIKVPSDQEILDAHNKYYADIKLDTNTGNTYIDGIYFEPTQTNKEALDY
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unk VIRT2310 Blast_submission tr Q46221 Q46221 CLOPE	 VMTYKKLRIYAVTPDNRELLVLSVN

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In case of problems, please read the online BLAST help.

If your question is not covered, please contact <helpdesk@expasy.org?

NCBI BLAST program reference [PMID:9254694]:

Altschul S.F., Madden T.L., Schäffer A.A., Zhang J., Zhang Z., Mille: Lipman D.J. Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res. 25:3389-3402(1997).

Query: 23 AA

Date run: 2005-11-30 16:03:15 UTC+0100 on sib-gml.unil.ch

Program: NCBI BLASTP 1.5.4-Paracel [2003-06-05]

Database: EXPASY/UniProtKB

2,618,771 sequences; 854,910,163 total letters

UniProt Knowledgebase Release 6.5 consists of:

UniProtKB/Swiss-Prot Release 48.5 of 22-Nov-2005: 199607 entries UniProtKB/TrEMBL Release 31.5 of 22-Nov-2005: 2406391 entries



List of potentially matching sequences

Send se	elected sec	quences to
Clustal \	V (multiple a	lignment) 🔽 Submit Query
Selec	£ up to	
☐ Incl	ude query	sequence
Db) AC	Description
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     75015 Paris Cedex 15, FRANCE
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for liota toxin clostridium per



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[Entry info] [Name and origin] [References] [Comments] [Cross-references] [Keywords] [Features] [Sequence] [Tools]

Note: most headings are clickable, even if they don't appear as links. They link to the user manual or other documents.

Entry information

Entry name

Q46221 CLOPE

Primary accession number

Entered in TrEMBL in

Q46221

Secondary accession numbers

None Release 01, November 1996

Sequence was last modified in

Release 01, November 1996

Annotations were last modified in

Release 24, June 2003

Name and origin of the protein

Protein name

lota toxin component lb [Precursor]

Synonyms

None None

Gene name From

Clostridium perfringens [TaxID: 1502]

Taxonomy

Bacteria; Firmicutes; Clostridia; Clostridiales; Clostridiace

Clostridium.

References

[1] NUCLEOTIDE SEQUENCE.

STRAIN=NCIB 10748;

PubMed=8225592 [NCBI, ExPASy, EBI, Israel, Japan]

Perelle S., Gibert M., Boquet P., Popoff M.R.;

"Characterization of Clostridium perfringens iota toxin genes and expression in Escherichi coli.":

Infect. Immun. 61:5147-5156(1993).

[2] NUCLEOTIDE SEQUENCE.

STRAIN=NCIB 10748:

Popoff M.R.:

Submitted (AUG-1995) to the EMBL/GenBank/DDBJ databases.

Comments

None

Cross-references

X73562; CAA51960.1; -;

[EMBL / GenBank / DDBJ]

[CoDingSequence] Genomic DNA.

PIR

EMBL

140862; 140862.

HSSP P13423; 1ACC. [HSSP ENTRY / PDB] GO:0005576; Cellular component: extracellular region (inferred from electro

annotation).

GO:0009405; Biological process: pathogenesis (inferred from electronic

annotation).

QuickGo view.

IPR003896; Anthrax toxinB.

InterPro IPR011658; PA14.

Graphical view of domain structure.

PF03495; Binary toxB; 1.

Pfam PF07691: PA14: 1.

Pfam graphical view of domain structure.

PRINTS PR01391; BINARYTOXINB.

ProDom [Domain structure / List of seq. sharing at least 1 domain]

HOGENOM [Family / Alignment / Tree]

ProtoMap Q46221. PRESAGE Q46221. ModBase Q46221.

SWISS-

2DPAGE Get region on 2D PAGE.

UniRef View cluster of proteins with at least 50% / 90% / 100% identity.

Keywords

Signal.

Features



Feature table viewer

KeyFromToLengthDescriptionSIGNAL34396Potential.

CHAIN 212 875 664 iota toxin component Ib.

Sequence information

Length: 875 AA [This is the length of the unprocessed precursor]

Molecular weight: 98469 Da [This is the MW of the unprocessed precursor]

CRC64: C9AE092CD3818921 is a checksum on the sequenc

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ScanProsite, MotifScan

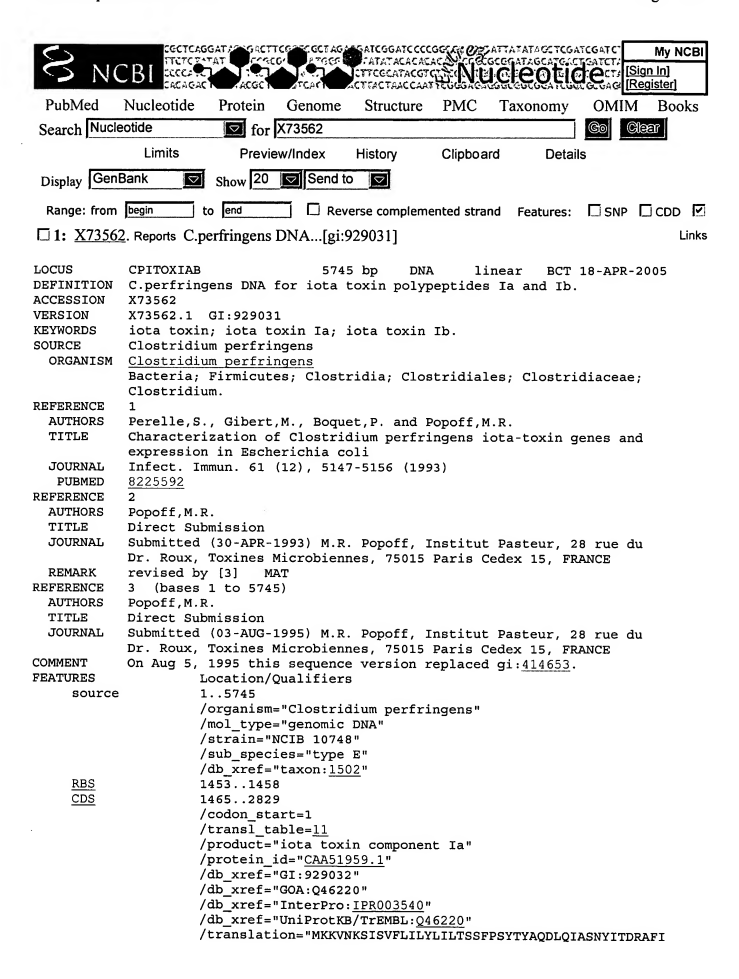


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General Information

Primary Accession # X73562

Accession #

X73562

Entry Name

EMBL:CPITOXIAB

Molecule Type

genomic DNA

Sequence Length

5745

Entry Division

PRO

Sequence Version

X73562.1

Creation Date

04-NOV-1993

Modification Date

18-APR-2005

Description

Description

C.perfringens DNA for iota toxin polypeptides Ia and Ib

Keywords

iota toxin; iota toxin Ia; iota toxin Ib.;

Organism

Clostridium perfringens

Organism

Classification

Bacteria; Firmicutes; Clostridia; Clostridiales; Clostridiaceae; Clostridium.

References

1. Perelle, S.; Gibert, M.; Boquet, P.; Popoff, M.R.;

Characterization of Clostridium perfringens iota toxin gene expression in Escherichia coli

Infect. Immun. 61(12):5147-5156 (1993)

Pubmed 8225592

- **2.** Popoff, M.R.; Submitted (30-APR-1993) to the EMBL/GenBank/DDBJ databases. Popoff, Institut Pasteur, 28 rue du Dr. Roux, Toxines Microbiennes, 75015 Paris (**FRANCE**
- 3. Popoff, M.R.; Submitted (03-AUG-1995) to the EMBL/GenBank/DDBJ databases Popoff, Institut Pasteur, 28 rue du Dr. Roux, Toxines Microbiennes, 75015 Paris (**FRANCE**

organism Clostridium perfringens

Position 1-5745

Additional Information

Features

Key		Location	Qualifier	Value
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			mol_type	genomic DNA

sub_species type E

strain NCIB 10748

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db xref GOA:Q46220

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product iota toxin component Ib

Sequence

Characteristics

Length: 5745 BP, A Count:2397, C Count:663, G Count:803, T Count:1882, Others Count:0

Sequence

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TITLE: Targeting antigens to the MHC class I processing pathway with an anthrax toxin fusion protein

Brief Summary Text (97):

In general, for cloning and expression of PA, the same methods as described for antigen-APABP can be used by one skilled in the art. Genes that encode wild type or mutated proteins can be cloned and expressed by methods known to those skilled in the art, as described above. For example, the gene encoding protein Ib of the Clostridium perfringens iota toxin can be cloned and expressed for use in the present invention according the methods described herein, or by methods known to those skilled in the art. The present invention uses an isolated nucleic acid in expression vector pYS5 that encodes the PA protein, as described in Example 2.

Other Reference Publication (4):

Sirard, Jean-Claude, et al. (1997) "A Recombinant Bacillus anthracis Strain Producing the Clostridium perfringens <u>Ib Component Induces Protection against Iota</u> Toxins", Infection and Immunity, 65(6): 2029-2033.

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Perelle 8., Gibert M., Boquet P., Popoff M.R.;
Forelle 8., Gibert M., Boquet P., Popoff M.R.;
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Forestle 1 Strokethola coli.;
Infect. Immun. 61:5147-5156(1993).
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Infect. Immun., 12 1993, 5147-5156, Vol 61, No. 12 Copyright © 1993, American Society for Microbiology

Characterization of Clostridium perfringens iota-toxin genes and expression in Escherichia coli [published erratum appears in Infect Immun 1995 Dec;63(12):4967]

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S Perelle, M Gibert, P Boquet and MR Popoff Laboratoire des Toxines Microbiennes, Institut Pasteur, Paris, France.

The iota toxin which is produced by Clostridium perfringens type E, is a binary toxin consisting of two independent polypeptides: Ia, which is an ADP-ribosyltransferase, and Ib, which is involved in the binding and internalization of the toxin into the cell. Two degenerate oligonucleotide probes deduced from partial amino acid sequence of each component of C. spiroforme toxin, which is closely related to the iota toxin, were used to clone three overlapping DNA fragments containing the iota-toxin genes from C. perfringens type E plasmid DNA. Two genes, in the same orientation, coding for Ia (387 amino acids) and Ib (875 amino acids) and separated by 243 noncoding nucleotides were identified. A predicted signal peptide was found for each component, and the secreted Ib displays two domains, the propeptide (172 amino acids) and the mature protein (664 amino acids). The Ia gene has

been expressed in Escherichia coli and C. perfringens, under the control of its own promoter. The recombinant polypeptide obtained was recognized by Ia antibodies and ADP-ribosylated actin. The expression of the Ib gene was obtained in E. coli harboring a recombinant plasmid encompassing the putative promoter upstream of the Ia gene and the Ia and Ib genes. Two residues which have been found to be involved in the NAD+ binding site of diphtheria and pseudomonas toxins are conserved in the predicted Ia sequence (Glu-14 and Trp-19). The predicted amino acid Ib sequence shows 33.9% identity with and 54.4% similarity to the protective antigen of the anthrax toxin complex. In particular, the central region of Ib, which contains a predicted transmembrane segment (Leu-292 to Ser-308), presents 45% identity with the corresponding protective antigen sequence which is involved in the translocation of the toxin across the cell membrane.

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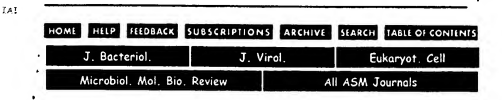
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